

## **STIC Biotechnology Systems Branch**

### **RAW SEQUENCE LISTING** **ERROR REPORT**

The Biotechnology Systems Branch of the Scientific and Technical Information Center (STIC) detected errors when processing the following computer readable form:

Application Serial Number: 10/564,484  
Source: IFWP  
Date Processed by STIC: 1/20/06

THE ATTACHED PRINTOUT EXPLAINS DETECTED ERRORS.

PLEASE FORWARD THIS INFORMATION TO THE APPLICANT BY EITHER:

- 1) INCLUDING A COPY OF THIS PRINTOUT IN YOUR NEXT COMMUNICATION TO THE APPLICANT, WITH A NOTICE TO COMPLY or,
- 2) TELEPHONING APPLICANT AND FAXING A COPY OF THIS PRINTOUT, WITH A NOTICE TO COMPLY

FOR CRF SUBMISSION AND PATENTIN SOFTWARE QUESTIONS, PLEASE CONTACT MARK SPENCER, TELEPHONE: 571-272-2510; FAX: 571-273-0221

TO REDUCE ERRORED SEQUENCE LISTINGS, PLEASE USE THE **CHECKER VERSION 4.4.0 PROGRAM**, ACCESSIBLE THROUGH THE U.S. PATENT AND TRADEMARK OFFICE WEBSITE. SEE BELOW FOR ADDRESS:

**<http://www.uspto.gov/web/offices/pac/checker/chkrnote.htm>**

Applicants submitting genetic sequence information electronically on diskette or CD-Rom should be aware that there is a possibility that the disk/CD-Rom may have been affected by treatment given to all incoming mail.

Please consider using alternate methods of submission for the disk/CD-Rom or replacement disk/CD-Rom.

Any reply including a sequence listing in electronic form should NOT be sent to the 20231 zip code address for the United States Patent and Trademark Office, and instead should be sent via the following to the indicated addresses:

1. EFS-Bio (<<http://www.uspto.gov/ebc/efs/downloads/documents.htm>> , EFS Submission User Manual - ePAVE)
2. U.S. Postal Service: Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450
3. Hand Carry, Federal Express, United Parcel Service, or other delivery service (EFFECTIVE 01/14/05):  
U.S. Patent and Trademark Office, Mail Stop Sequence, Customer Window, Randolph Building, 401 Dulany Street, Alexandria, VA 22314

Revised 01/10/06

### Raw Sequence Listing Error Summary

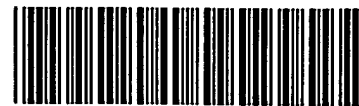
**ERROR DETECTED**

### SUGGESTED CORRECTION

SERIAL NUMBER: 10/564,484

**ATTN: NEW RULES CASES: PLEASE DISREGARD ENGLISH "ALPHA" HEADERS, WHICH WERE INSERTED BY PTO SOFTWARE**

- |    |  |  |
|----|--|--|
| 1  | _____ Wrapped Nucleics<br>Wrapped Aminos | The number/text at the end of each line “wrapped” down to the next line. This may occur if your file was retrieved in a word processor <b>after</b> creating it. Please adjust your right margin to .3; this will prevent “wrapping.”  |
| 2  | _____ Invalid Line Length                | The rules require that a line <b>not exceed</b> 72 characters in length. This includes white spaces.   |
| 3  | _____ Misaligned Amino<br>Numbering      | The numbering under each 5 <sup>th</sup> amino acid is misaligned. Do <b>not</b> use tab codes between numbers; use <b>space characters</b> , instead.   |
| 4  | _____ Non-ASCII                          | The submitted file was <b>not</b> saved in ASCII(DOS) text, as <b>required</b> by the Sequence Rules. <b>Please ensure your subsequent submission is saved in ASCII text.</b>  |
| 5  | _____ Variable Length                    | Sequence(s)_____ contain n’s or Xaa’s representing more than one residue. <b>Per Sequence Rules, each n or Xaa can only represent a single residue.</b> Please present the <b>maximum</b> number of each residue having variable length and indicate in the <220>-<223> section that some may be missing.  |
| 6  | _____ PatentIn 2.0<br>“bug”              | A “bug” in PatentIn version 2.0 has caused the <220>-<223> section to be missing from amino acid sequences(s) _____. Normally, PatentIn would automatically generate this section from the previously coded nucleic acid sequence. Please manually copy the relevant <220>-<223> section to the subsequent amino acid sequence. <b>This applies to the mandatory &lt;220&gt;-&lt;223&gt; sections for Artificial or Unknown sequences.</b>   |
| 7  | _____ Skipped Sequences<br>(OLD RULES)   | Sequence(s) _____ missing. If intentional, please insert the following lines for <b>each</b> skipped sequence:<br>(2) INFORMATION FOR SEQ ID NO:X: (insert SEQ ID NO where “X” is shown)<br>(i) SEQUENCE CHARACTERISTICS: (Do not insert any subheadings under this heading)<br>(xi) SEQUENCE DESCRIPTION:SEQ ID NO:X: (insert SEQ ID NO where “X” is shown)<br>This sequence is intentionally skipped<br><br>Please also adjust the “(ii) NUMBER OF SEQUENCES:” response to <b>include</b> the skipped sequences. |
| 8  | _____ Skipped Sequences<br>(NEW RULES)   | Sequence(s) _____ missing. If <b>intentional</b> , please insert the following lines for <b>each</b> skipped sequence.<br><210> sequence id number<br><400> sequence id number<br>000  |
| 9  | _____ Use of n’s or Xaa’s<br>(NEW RULES) | Use of n’s and/or Xaa’s have been detected in the Sequence Listing.<br>Per 1.823 of Sequence Rules, use of <220>-<223> is <b>MANDATORY</b> if n’s or Xaa’s are present.<br>In <220> to <223> section, please explain location of <b>n</b> or <b>Xaa</b> , and which residue <b>n</b> or <b>Xaa</b> represents.   |
| 10 | _____ Invalid <213><br>Response          | Per 1.823 of Sequence Rules, the only <b>valid</b> <213> responses are: Unknown, Artificial Sequence, or scientific name (Genus/species). <220>-<223> section is <b>required</b> when <213> response is Unknown or is Artificial Sequence  |
| 11 | _____ Use of <220>                       | Sequence(s) _____ missing the <220> “Feature” and associated numeric identifiers and responses. Use of <220> to <223> is <b>MANDATORY</b> if <213> “Organism” response is “Artificial Sequence” or “Unknown.” Please explain source of genetic material in <220> to <223> section.<br>(See “Federal Register,” 06/01/1998, Vol. 63, No. 104, pp. 29631-32) (Sec. 1.823 of Sequence Rules)  |
| 12 | _____ PatentIn 2.0<br>“bug”              | Please do not use “Copy to Disk” function of PatentIn version 2.0. This causes a corrupted file, resulting in missing mandatory numeric identifiers and responses (as indicated on raw sequence listing). Instead, please use “File Manager” or any other manual means to copy file to floppy disk.  |
| 13 | _____ Misuse of n/Xaa                    | “n” can <b>only</b> represent a single nucleotide; “Xaa” can <b>only</b> represent a single amino acid   |



IFWP

## RAW SEQUENCE LISTING

DATE: 01/20/2006

PATENT APPLICATION: US/10/564,484

TIME: 12:30:46

Input Set : N:\SMITH\pto.ts20..txt

Output Set: N:\CRF4\01202006\J564484.raw

3 <110> APPLICANT: Japan Science and Technology Corporation  
 5 <120> TITLE OF INVENTION: Diagnosis method of endometriosis  
 7 <130> FILE REFERENCE: 03-F-104PCT  
 C--> 9 <140> CURRENT APPLICATION NUMBER: US/10/564,484  
 C--> 9 <141> CURRENT FILING DATE: 2006-01-13  
 9 <150> PRIOR APPLICATION NUMBER: JP2003-196459  
 10 <151> PRIOR FILING DATE: 2003-07-14  
 12 <160> NUMBER OF SEQ ID NOS: 10  
 14 <170> SOFTWARE: PatentIn version 3.1

*see item 4  
on Error Summary  
Sheet*

**Does Not Comply  
Corrected Diskette Needed**

## ERRORED SEQUENCES

177 <210> SEQ ID NO: 8  
 178 <211> LENGTH: 22 *29 shown below*  
 179 <212> TYPE: DNA  
 180 <213> ORGANISM: Artificial  
 182 <220> FEATURE:  
 183 <223> OTHER INFORMATION: Description of Artificial sequence: Synthetic  
 oligonucleotide  
 185 <400> SEQUENCE: 8  
 E--> 186 ttgaattctt aacatttctc catctctaa 29  
 201 <210> SEQ ID NO: 10  
 202 <211> LENGTH: 28  
 203 <212> TYPE: DNA  
 204 <213> ORGANISM: Artificial  
 206 <220> FEATURE:  
 207 <223> OTHER INFORMATION: Description of Artificial sequence: Synthetic  
 oligonucleotide  
 209 <400> SEQUENCE: 10  
 210 ggccgaattc agatccaaaa taattgcc 28  
 E--> 211 *1/5 delete*

**VERIFICATION SUMMARY**

PATENT APPLICATION: US/10/564,484

DATE: 01/20/2006

TIME: 12:30:47

Input Set : N:\SMITH\pto.ts20..txt

Output Set: N:\CRF4\01202006\J564484.raw

L:9 M:270 C: Current Application Number differs, Replaced Current Application No  
L:9 M:271 C: Current Filing Date differs, Replaced Current Filing Date  
L:26 M:258 W: Mandatory Feature missing, <223> Blank for SEQ#:1,Line#:24  
L:186 M:252 E: No. of Seq. differs, <211> LENGTH:Input:22 Found:29 SEQ:8  
L:211 M:254 E: No. of Bases conflict, LENGTH:Input:5 Counted:29 SEQ:10  
L:211 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:2  
L:211 M:252 E: No. of Seq. differs, <211> LENGTH:Input:28 Found:29 SEQ:10